Entry of *Trypanosoma (Schizotrypanum) dionisii* to Macrophages *in vitro* and its Subsequent Fate Therein

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Observations by phase contrast, fluorescence and electron microscopy showed that epimastigotes of *Trypanosoma (Schizotrypanum) dionisii* (grown *in vitro*) were phagocytosed posterior end first by mouse peritoneal macrophages *in vitro*. Many were subsequently digested as a result of phagosome-lysosome fusion but others survived by apparently inhibiting this fusion and/or escaping from the phagosome into the host cell's cytoplasm. These survivors replicated as amastigotes. Long trypomastigotes, separated from populations grown *in vitro* by passage down a column of glass beads (with or without prior exposure to guinea-pig serum), were phagocytosed by either pole and all were subsequently digested.

**INTRODUCTION**

*Trypanosoma dionisii* is phagocytosed by mouse peritoneal macrophages *in vitro* and some organisms survive to replicate intracellularly (Baker & Liston, 1978). This paper describes the process of endocytosis and the parasites' subsequent fate. Some preliminary observations have been published (Liston, 1975; Baker et al., 1977).

**METHODS**

Trypanosomes. *Trypanosoma (Schizotrypanum) dionisii* Bettencourt & Franca, 1905 was maintained in L4NHS medium at 28 °C. Unless otherwise specified, clone no. 3 of stock p3 was used; in some experiments uncloned stock p3, or the similar stock p2, were employed. Details of isolation, maintenance and cloning are given by Baker et al. (1976).

Macrophages. Peritoneal macrophages were collected from unstimulated male Parke's surgically derived [P(SD)] mice (Baker & Liston, 1978). For some electron microscopical observations, donor mice had been injected intradermally with 10⁸ *T. dionisii* p3 emulsified in Freund's Incomplete Adjuvant 28 d previously.

Light microscopy. Portions (1.5 ml) of a macrophage suspension (3 × 10⁴ ml⁻¹ in medium 199; Baker & Liston, 1978) were incubated at 37 °C in Wild Selecta tissue culture observation chambers (Leitz, Luton, LU1 3HP) for 24 h. The medium was then replaced by 199 containing 20 % (v/v) Wellcome heat-inactivated calf serum number 1 (WCS1) and 0-9 × 10⁴ to 2-1 × 10⁴ ml⁻¹ epimastigotes or trypomastigotes of clone 3 or stock p2 prepared according to Baker et al. (1976) (sometimes omitting the initial treatment with guinea-pig serum). Cultures were kept at 35 or 37 ± 0.1 °C by a thermistor-controlled fan heater (Klein & Law, 1976), examined by phase contrast microscopy and photographed by electronic flash illumination on Ilford Pan F film (ASA25).

Some cultures were supravitally stained with acridine orange 21667 (G. T. Gurr) in medium 199 (10 μg ml⁻¹) for 25 to 60 (usually 30) min at 37 °C (D'ArCY Hart & Young, 1975), either 0 to 90 min before or 24 to 72 h after addition of trypanosomes. Cultures were then rinsed with medium 199 before incubation in medium 199 with serum (37 °C, 1 to 4 h) to allow localization of stain within lysosomes. Cultures were

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examined by phase contrast and dark-field fluorescence microscopy (λ max. ~ 400 nm, Leitz Heine condenser) and photographed on Kodak Ektachrome film (ASA160).

Electron microscopy. (i) Monolayer cultures. Plastic tissue culture flasks (Falcon 25 cm²; Becton, Dickinson & Co., Wembley, HA9 OPS) contained 4x10⁶ macrophages in 8 ml medium 199 with 20% WCS1 or 16% WCS1 plus 4% heat-inactivated calf serum (Gibco Bio-Cult, Paisley, PA3 4EP) to ensure better cell adherence: 24 h later 2x10⁶ T. dionisii p3 (uncloned) were added. After 5 d at 32 °C or 3 d at 35 °C, cells were harvested by treatment (5 min, 37 °C) with 0.25% (w/v) trypsin (Difco, 1:2500) in phosphate-buffered saline (PBS; Dulbecco’s solution A, Oxoid), washed twice in PBS, centrifuged (2000 g, 5 min) and fixed at 4 °C either for 1 h in osmium tetroxide/glutaraldehyde solution (Hirsch & Fedorko, 1968) in 0.2 m-cacodylate buffer or for 2 h in 5% (v/v) glutaraldehyde in 0.1 m-Sorenson’s phosphate buffer (SB, pH 7.4). They were then washed (4 °C, 400 g, 5 min) in 0.2 m-cacodylate buffer plus 5% (w/v) sucrose or in SB and centrifuged (50 °C, 600 g, 5 min) in molten 1% (w/v) Ionagar no. 1 (Oxoid). Cubes (about 1 mm³) were postfixed for 2 h in 1% (w/v) osmium tetroxide in SB (if initially fixed with glutaraldehyde only), washed in SB, dehydrated in ethanol and embedded in Epon via propylene oxide. Sections were stained with 2% (w/v) aqueous uranyl acetate and aqueous lead citrate (Reynolds, 1963) for 2 min each.

(ii) Suspension cultures. Washed T. dionisii (as above) were added to macrophages (7 trypanosomes per macrophage) suspended in medium 199 plus 20% WCS1 in 2 ml screw-capped plastic ampoules (no. 506; Sterilin, Teddington, TW11 8QZ). After 10 or 20 min at 37 °C with occasional shaking, cultures were centrifuged (2000 g, 400 g, 5 min) and fixed in 2.5% glutaraldehyde in SB, centrifuged in agar, postfixed (1 or 2 h), dehydrated and embedded as described above (but using Taab resin; Taab Laboratories, Reading, Berks).

(iii) Lysosomal labelling. (a) Monolayer cultures were treated, 24 h after initiation, with ferritin (Taab) in medium 199 (0.5 mg ml⁻¹; 4 h, 37 °C; Jones & Hirsch, 1972), washed and then incubated in medium 199 with 20% serum (4 h, 37 °C). Clone 3 (mainly epimastigotes) were added at 10 trypanosomes per macrophage. After 1 h at 37 °C, cultures were fixed in 2% (v/v) glutaraldehyde in SB (1 h, 4 °C); cells were scraped off the substrate (Alexander & Vickerman, 1975), centrifuged in agar, postfixed and embedded as described above. (b) Cultures of 2x10⁶ macrophages in 2 ml medium 199 in 28 ml glass Universal bottles were incubated, about 18 h after initiation, in 0.2% (w/v) saccharated iron oxide in medium 199 (3 h, 37 °C; Alexander & Vickerman, 1975), washed and incubated in medium 199 plus 10% WCS1 for 1 h. Then 1x10⁶ epimastigotes of clone 3 were added. After 18 h, cultures were fixed and prepared for electron microscopy as described by Alexander & Vickerman (1975).

RESULTS

Attachment and entry to macrophages

Early experiments were at 32 or 35 °C, but most were at 37 °C. No difference was observed between events occurring at the different temperatures.

Immediately after their introduction, epimastigotes became attached to macrophages by their posterior tip. Entry followed during the next 10 min, apparently by phagocytosis (Figs 1 and 2) (Baker & Liston, 1978). Trypomastigotes, however, became attached by either their anterior or posterior ends, and entered (again apparently by phagocytosis) either end first (depending which end had become attached to the cell). After entry, both forms became coiled up in parasitophorous vacuoles and remained motile, the trypomastigotes in particular moving like coiled springs (Fig. 3). Many endocytosed epimastigotes and all trypomastigotes were digested within 24 to 48 h, but some epimastigotes survived, became rounded and

Figs 1 to 3. Serial phase contrast photomicrographs of an epimastigote (p) of T. dionisii p3 being phagocytosed by a mouse peritoneal macrophage at 35 °C in vitro: Fig. 1, 0 min; Fig. 2, 7 min; Fig. 3, 10 min. Bar markers represent 10 μm.

Fig. 4. Phase contrast photomicrograph of a macrophage after exposure in vitro to T. dionisii p3 at 37 °C for 5 d. Some intracellular amastigotes (a) appear to be in vacuoles and others (a') free in the cytoplasm; some have become vacuolated (av). Bar marker represents 10 μm.

Figs 5 and 6. Paired phase contrast (a) and fluorescence (b) photomicrographs of macrophages exposed to T. dionisii p3 epimastigotes at 37 °C and labelled in vitro with acridine orange either 90 min before (Fig. 5) or 48 h after (Fig. 6) exposure to parasites. Some live parasites (p) are adjacent to host-cell secondary lysosomes (hs). A large phagolysosome (Fig. 5, hpl) contains a vacuolated non-motile parasite (p'). In Fig. 6, one cell containing many intact amastigotes (a) entirely lacks fluorescence label. Bar markers represent 10 μm.
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subsequently replicated by binary fission (Fig. 4). Epimastigotes also multiplied extracellularly in the supernatant medium, and continued to enter macrophages throughout the observation period (96 h). Consequently, rolled-up epimastigotes and amastigotes were seen throughout this period; later, some of them were not surrounded by obvious vacuoles (compare electron microscopic observations reported below).

**Fluorescence microscopy**

Acridine orange in lysosomes of macrophages and parasites fluoresced bright orange-red under blue-violet light. More lysosomes were seen in macrophages that were well spread after several days in culture. Some macrophages which appeared rounded under phase contrast illumination were packed with lysosomes, presumably indicating autolysis. Lysosomes were also seen at the posterior tip of parasites, epimastigotes having up to four or five and trypomastigotes only one large one. Trypanosome kinetoplasts and nuclei of parasites and macrophages fluoresced green.

Addition of trypanosomes immediately after applying acridine orange to cultures was unsatisfactory, resulting in a green halo around living attached and intracellular parasites. When the dye was allowed to localize in the lysosomes of macrophages for 90 min before 3 h exposure to trypanosomes, living intracellular parasites, whether in large phagosomes or not, were not associated with dye. Large lysosomes could be seen next to living parasites but fusion with the parasitophorous vacuoles had not occurred. Trypanosomes obviously undergoing digestion were surrounded by large fluorescing phagolysosomes (Fig. 5). If cultures were infected 48 h before exposure to acridine orange, cells in which much digestion was evident were heavily labelled. Again living parasites not in large parasitophorous vacuoles had no associated dye. One cell harbouring several apparently viable amastigotes contained no detectable fluorochrome (Fig. 6). Macrophages in cultures infected 72 h before addition of dye contained numerous fluorescing lysosomes, which surrounded some of the living dividing intracellular parasites but had not fused with the latter's parasitophorous vacuoles. Some viable parasites were completely unassociated with lysosomes; yet others, and all digested parasites, were contained in large fluorescing vacuoles, indicating phagosome-lysosome fusion.

Macrophages in cultures with pure populations of trypomastigotes invariably showed fusion of lysosomes with parasitophorous vacuoles, often even while the parasite was being ingested. This correlated with the observed occurrence in Giemsa-stained control preparations of their complete digestion within 24 to 48 h.

**Electron microscopy**

In monolayer cultures, after phagocytosis by macrophages (Fig. 7), rolled-up epimastigotes or amastigotes were often seen surrounded by a closely applied membrane (Fig. 8) or apparently free in the cytoplasm of macrophages (Fig. 9). In cells incubated at 32 °C, they

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Figs 7 to 13. Electron micrographs of thin sections of mouse peritoneal macrophages cultured in vitro with *T. dionisi* i3 (Figs 10, 11) or clone 3 (Figs 7, 8, 9, 12, 13). All except Fig. 11 were from monolayer cultures; Figs 7, 8, 9, macrophages labelled with saccharated iron oxide and exposed for 18 h to epimastigotes at 37 °C; Fig. 10, macrophage from an immunized mouse after exposure to *T. dionisi* at 32 °C for 5 d; Fig. 11, macrophage from a suspension culture 20 min after addition of *T. dionisi* at 37 °C; Figs 12, 13, macrophages labelled with ferritin and exposed for 1 h to *T. dionisi* at 37 °C. Bar markers represent 1 μm.

Fig. 7. Dividing epimastigote (p) apparently undergoing phagocytosis posterior end foremost.

Fig. 8. Parasite (p) in closely applied vacuole (hpv) with no label, alongside a similar vacuole containing label but no parasite, presumably a phagolysosome (hpl).

Fig. 9. Intact parasite (p) apparently free in cytoplasm alongside a vacuole containing a digested parasite (p') with associated label, indicating phagosome-lysosome fusion.
For general details, see legend on p. 256.

Fig. 10. Intracellular parasites, some dividing (dp), with associated host-cell cytolysosomes (hcy).

Fig. 11. Rolled-up parasite [with nucleus (pn), kinetoplast (k), flagellum (fm) and mitochondrion (pm)] surrounded by host-cell mitochondria (hcm) and endoplasmic reticulum (her).

Fig. 12. Labelled secondary lysosomes (hsl) and endoplasmic reticulum (her); a parasite (p) undergoing phagocytosis has no associated label.

Fig. 13. Secondary lysosomes (hsl) and larger vacuoles, presumably phagolysosomes (hpl) containing parasite material (fm) and associated label. An intact parasite (p) is in a parasitophorous vacuole without label.
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were frequently next to membrane-bound vesicles containing partially digested host-cell organelles, presumably cytolysosomes or autophagosomes (Fig. 10), the intervening membrane often being thick and osmiophilic; such appearances were less common in cells incubated at 35 °C, though no other consequence of the temperature difference was noticed. Dividing parasites, probably amastigotes, were seen (Fig. 10). No obvious difference was noted between infected macrophages from normal or immune mice. In suspension cultures only, some epimastigotes were attached to macrophages by their flagella and some parasitophorous vacuoles were surrounded by host-cell mitochondria and endoplasmic reticulum (Fig. 11).

Macrophages treated with ferritin before being exposed to trypanosomes for 1 h contained the label in their endoplasmic reticulum, primary and secondary lysosomes, and presumed autophagosomes (Fig. 12). Parasites undergoing phagocytosis had no associated ferritin (Fig. 12), nor did those intracellular forms surrounded by a closely applied membrane, but larger vacuoles, presumably phagolysosomes, contained parasites and ferritin (Fig. 13).

Macrophages treated with saccharated iron oxide and subsequently exposed to trypanosomes for 18 h possessed large labelled vacuoles (some also containing parasite debris), presumably phagolysosomes (Fig. 9). An apparently viable amastigote with no surrounding host membrane had no associated label, while an adjacent vacuole contained a digested parasite and label, indicating phagosome-lysosome fusion (Fig. 9); another intact parasite surrounded by a closely applied membrane was unlabelled, while a nearby large vacuole contained the oxide (Fig. 8).

**Discussion**

The inference from direct observations that parasites enter macrophages passively by phagocytosis, after attachment to the host-cell membrane, is supported by the inhibition of entry into cells treated with cytochalasin B or trypsin (Baker & Liston, 1978). In monolayer cultures this attachment was almost invariably by the posterior pole of epimastigotes and by either end of trytomastigotes. Most previous workers also reported entry of trypanosomatids posterior end first (see, for example, Akiyama & Haight, 1971; Nogueira & Cohn, 1976); in contrast, Miller & Twohy (1967) reported that phagocytosis by macrophages of *Leishmania donovani* promastigotes occurred flagellar end first.

The observed failure of any tryptomastigotes to survive within macrophages contrasts with the observation by Baker et al. (1972) that higher infection rates resulted in HeLa cells infected with populations containing higher proportions of these forms. Nogueira & Cohn (1976) found that tryptomastigotes of *T. cruzi*, separated from *in vitro* cultures by treatment with guinea-pig serum followed by albumen gradient centrifugation, could initiate an intracellular cycle in mouse peritoneal macrophages *in vitro*, whereas epimastigotes could not. Perhaps passage down a column of glass beads, part of our preparatory procedure (Baker et al., 1976), impaired in some way the tryptomastigotes' subsequent resistance to intracellular digestion, though they remained fully motile. The effect cannot have been due to exposure to guinea-pig serum, since inocula of tryptomastigotes of stock p2 prepared without this treatment failed to survive within macrophages; also, Nogueira & Cohn's (1976) preparations had been treated with guinea-pig serum.

The observations on preparations stained with acridine orange suggest that host-cell lysosomes may fuse very rapidly with parasitophorous vacuoles containing tryptomastigotes or epimastigotes — even before completion of phagocytosis of the former and at least within 1 h of ingestion of the latter. It appeared that all parasites within the resulting phagolysosomes were subsequently digested. However, some intracellular epimastigotes survived, either by escaping from the parasitophorous vacuoles into the host-cell cytoplasm or by apparently inhibiting lysosomal fusion with the vacuoles (which remained small and barely detectable by light microscopy) as reported by Khavkin & Freidlin (1977). The presence of acridine orange in vacuoles containing apparently viable parasites in macrophages labelled
72 h previously could have been due to recent endocytosis of surviving replicating extracellular parasites and need not imply resistance to digestion for the full 72 h. Electron microscopy supported this view, by showing that all morphologically intact and replicating parasites were either surrounded by a closely apposed host-cell membrane, enclosing no lysosomal marker, or were apparently free in the macrophage's cytoplasm. Resistance of *T. cruzi* to digestion has been ascribed to its escape from the phagosome (see, for example, Nogueira & Cohn, 1976; Kress et al., 1975, 1977; Milder et al., 1976, 1977), in contrast to *Leishmania* spp., which seem able to survive even in the presence of lysosomal contents within phagolysosomes (Alexander & Vickerman, 1975; Chang & Dwyer, 1976, 1978; Lewis & Peters, 1977), and *Toxoplasma gondii* which was described as being 'walled off' from host-cell lysosomes by the macrophage's mitochondria and endoplasmic reticulum (Jones & Hirsch, 1972). The significance of this 'walling off', also seen in some of our material, is not clear; it appeared to be an exceptional occurrence probably resulting from mechanical displacement of the host-cell organelles by the phagocytic vacuole.

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**REFERENCES**


